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## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

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**(54) Title: MATERIALS AND METHODS RELATING TO DISEASE DIAGNOSIS**  
**(54) Titre: MATERIAUX ET PROCEDES SE RAPPORTANT AU DIAGNOSTIC DE MALADIE**

**(57) Abstract**

The invention provides materials and methods relating to disease diagnosis. In particular, the invention provides a method of diagnosing diseases, such as cancers, by comparing specific patterns of gene expression characteristic of the disease at a nucleic acid or protein level. The invention provides novel methods for analysing the expression profiles characteristic of diseased cells, in order to determine specific diagnostic markers. Such determined diagnostic markers may be stored on, for example a database, and used in the diagnosis of diseases such as cancer.

**(57) Abrégé**

La présente invention concerne des matériaux et des procédés se rapportant au diagnostic de maladie. Elle concerne en particulier un procédé de diagnostic de maladies, telles que des cancers, qui consiste à comparer des formes spécifiques d'expression génique caractéristique de la maladie à un niveau d'acide nucléique ou de protéine. Elle concerne aussi des procédés d'analyse des profils d'expression caractéristique de cellules malades, de manière à déterminer des marqueurs spécifiques de diagnostic. Ces marqueurs déterminés peuvent être conservés, par exemple dans une base de données, et utilisés dans le diagnostic de maladies telles que le cancer.

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(54) Title: MATERIALS AND METHODS RELATING TO DISEASE DIAGNOSIS

(57) Abstract

The invention provides materials and methods relating to disease diagnosis. In particular, the invention provides a method of diagnosing diseases, such as cancers, by comparing specific patterns of gene expression characteristic of the disease at a nucleic acid or protein level. The invention provides novel methods for analysing the expression profiles characteristic of diseased cells, in order to determine specific diagnostic markers. Such determined diagnostic markers may be stored on, for example a database, and used in the diagnosis of diseases such as cancer.

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Materials and Methods Relating to Disease Diagnosis

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Field of the Invention

The present invention concerns materials and methods relating to disease diagnosis. Particularly, but not exclusively, the invention relates to methods of diagnosing tumours, by comparing specific patterns of gene expression at a nucleic acid or protein level using expressed nucleic acid, e.g. mRNA or cellular proteins associated with the tumour.

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Background of the Invention

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The major characteristics that differentiate malignant tumours from benign ones are their properties of invasiveness and spread. Malignant tumours do not remain localised and encapsulated: they invade surrounding tissues, get into the body's circulatory system, and set up areas of proliferation away from the site of their original appearance. When tumour cells spread and engender secondary areas of growth, the process is called metastasis; malignant cells having the ability to metastasize.

The earliest stages of malignant tumours are hard to identify and pathologists are rarely sure how or where a malignancy began. The cells of malignant tumours have a tendency to lose differentiated traits and therefore it can be difficult to determine the primary origin of the

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cells following metastasis.

A concern with the histopathologic assessment of neoplasias (tumour growth) is that tumour classification is based on subjective evaluation (1, 2). Immunostaining can be used to determine the expression of various diagnostic markers and may increase reproducibility.

Ovarian cancer is an example of a disease where the diagnostic difficulties are considerable (3). Epithelial neoplasias of ovarian cancers are classified into benign, borderline and malignant tumours. Borderline tumours are often difficult to diagnose, and it is not known if most of these tumours represent intermediate steps in tumour progression or whether these tumours should be considered as a separate group (4). Relative survival decreases with increasing tumour stage or grade. Five-year survival is considerably lower for women with carcinoma (38%) than for women with borderline carcinoma (95%).

#### Summary of the Invention

The present inventors have appreciated that carrying out routine tumour diagnosis in an accurate and objective manner is very difficult. The process is preoperatively dependent on an experienced cytologist and/or postoperatively dependent on an experienced pathologist, and is at present based on morphological judgements.

Further, the primary tumour source can be difficult to determine which may lead to miss-diagnosis and

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inappropriate treatment regime. Therefore, the present inventors have realised that there is a need for a diagnostic tool that can perform preoperative diagnosis objectively. Such a tool should help to reduce the number of patients undergoing unnecessary and expensive therapy.

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Multivariate analysis of the expression of a series of diagnostic markers is one approach to diagnostic problems. If a sufficiently large data set is collected, it may be possible to recognize patterns of expression in different histological groups. Goldschmidt et al. (5) showed that multivariate analysis of 47 histological variables generated by computer-assisted microscope analysis facilitated classification of adipose tumours. Similarly, multivariate analysis of RNA expression data has been used to discriminate between fibroblast subtypes (6).

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One approach to obtain a large data set is to use high resolution two-dimensional polyacrylamide gel electrophoresis (2-DE). This technique is able to resolve more than one thousand polypeptides on a single gel. The pattern can be analysed by computer software such as PDQUEST and MELANIE II (7, 8). This approach has been previously used for the classification of lung tumour cell lines (9).

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An alternative approach to obtaining a large data set is to use micro-array technology. Nucleic acid

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sequence characteristic of nucleic acid sequences expressed in certain cell types, e.g. mRNA or cDNA, can be analysed in this way. There is an increasing tendency towards miniaturisation of assays which use binding members (such as antibodies or nucleic acid sequences).

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For example, the binding members may be immobilised in small discrete locations (microspots) and/or as arrays (micro-array technology) on solid supports or on diagnostic chips. These approaches can be particularly valuable as they can provide great sensitivity (particularly through the use of fluorescent labelled reagents), require only very small amounts of biological sample from individuals being tested and allow a variety of separate assays to be carried out simultaneously.

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Examples of techniques enabling miniaturised technology are provided in WO84/01031, WO88/1058, WO89/01157, WO93/8472, WO95/18376, WO95/18377, WO95/24649 AND EP-A-0373203.

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Early research by Fedor et al established that silicon could serve as a substrate onto which organic molecules such as DNA could be synthesized. The process now commercialised by Affymetrix Inc. Santa Clara, California, involves the use of serial photolithographic steps to build oligonucleotides *in situ* at a specific addressable position on the chip.

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The strategy of addressing specific nucleic acid sequences synthesized off chip, then hybridized to a

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particular location on a chip by electrical attraction to  
a charged microelectrode has been developed by Nanogen  
10 Inc. Variation on the theme of microaddressable arrays  
has recently led to the evaluation of chips for sequence  
15 analysis of uncharaterised genetic material, mutational  
analysis of a known gene locus, and for the evaluation of  
a particular cell or tissue's profile of gene expression  
20 for the whole complement of the human DNA sequence. These  
methodologies typically relay on the use of laser  
25 activated fluorescence of addressable signals on a  
microchip.

Thus, at its most general, the present invention  
provides materials and methods for, firstly obtaining a  
30 number of protein or nucleic acid expression profiles  
15 characteristic for disease states of different origins or  
different stages of development or malignancy; secondly,  
analysing said expression profiles in order to determine  
35 specific diagnostic markers; and thirdly, diagnosing the  
presence of a disease, e.g. tumour, the type of disease  
20 or the stage of development of said disease e.g. tumour  
malignancy by comparison of its protein or nucleic acid  
40 expression profile with those previously obtained to  
determine using the specified diagnostic markers.

Thus, the present invention primarily relates to a  
45 method of obtaining gene expression profiles in order to  
25 determine diagnostic markers characteristic of a selected  
disease type or stage of development of a disease

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comprising

10 (1) obtaining cells from a sample of said disease  
tissue;

15 (2) disrupting cells to expose the cellular products  
characteristic of gene expression;

20 (3) separating said cellular products according to  
their characteristic properties on a substrate; and

25 (4) carrying out computer-assisted multivariate  
analysis of the substrate to quantify and characterise  
10 the cellular product distribution on the substrate to  
identify specific diagnostic markers characteristic of  
25 said disease.

30 Depending on the cell type, different genes are  
expressed or are expressed at different levels or  
35 frequency. These differences in gene expression may be  
used to characterise the type of cell. The cellular  
products that reflect the differences in gene expression  
are those products produced downstream of the nucleic  
acid transcription and translation process, e.g. mRNA or  
40 the expressed protein itself. These cellular products may  
then be separated according to their own characteristic  
properties, e.g. size, charge or sequence.

45 In a preferred embodiment of the invention, the  
cellular products are expressed proteins which may be  
50 separated according to their size on a electrophoresis  
gel, preferably a two dimensional electrophoresis gel.

55 Alternatively, the cellular products may be

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separated according to their characteristic properties using a substrate comprising specific binding members, for example, antibodies or oligonucleotides. As mentioned above, this is conveniently done by using a micro-array.

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5 In such a situation, it is preferable to label the cellular products, e.g. radioactively or fluorescently or enzymatically, to assist in the computer-assisted multivariate analysis.

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Therefore, in a first aspect, the present invention provides a method of obtaining protein expression profiles in order to determine diagnostic markers characteristic of selected disease types or stages of disease development comprising

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30 (1) obtaining cells from a sample of said disease type;

15 (2) disrupting cells to expose the cellular proteins contained therein;

35 (3) separating said cellular proteins using a two-dimensional electrophoresis gel; and

20 (4) carrying out computer-assisted multivariate analysis of the two-dimensional electrophoresis gel to 40 quantify and characterise the protein distribution on the gel to identify specific diagnostic markers characteristic of said disease.

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25 In order to carry out the analysis as outline in 50 step (4), quantitative and qualitative data from the two-dimensional electrophoresis gel is firstly obtained.

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Thus, step (4) may require carrying out multivariate analysis of the quantitative and qualitative data from the two-dimensional gel to characterise the protein expression profile and identify specific diagnostic markers characteristic of said disease.

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In an alternative first aspect of the present invention, there is provided a method of obtaining gene expression profiles in order to determine diagnostic markers characteristic of selected disease types or stages of disease development, said method comprising

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(1) obtaining cells from a sample of said disease

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type

(2) disrupting cells to obtain the expressed nucleic acid contained therein;

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(3) separating said expressed nucleic acid using a micro-array; and

(4) carrying out computer-assisted multivariate analysis of the micro-array to quantify and characterise the expressed nucleic acid on the micro-array to identify specific diagnostic markers.

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The expressed nucleic acid is preferably mRNA which may be obtained from the cells by standard molecular techniques known to the skilled person, for example see Sambrook, Fritsch and Maniatis, "Molecular Cloning, A Laboratory Manual", Cold Spring Harbor Laboratory Press, 1989, and Ausubel et al, Short Protocols in Molecular Biology, John Wiley and Sons, 1992). Alternatively, cDNA

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may be created from the expressed mRNA by reverse transcription before separation and analysing on the micro-array. Micro-array technologies use oligonucleotides (representing thousands of different genes) bound to given positions on various substrate. Total mRNA is purified from a cell/tissue sample and cDNA is produced by reverse transcriptase. Various steps (e.g. in vitro transcription using biotinylated nucleotides) may then be added before hybridisation and visualisation depending on the specific type of micro-array technology used (e.g. Affymetrix chips, Clontech membranes). The final read-out is a signal that is proportional to the quantity of a given expressed gene.

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The present inventors have discovered that proteins are differently expressed or differentially regulated between various malignant tumours and benign tumours.

Therefore, the inventors believe that the present invention will have particular utility in relation to the diagnosis of tumours. Although the following description of the invention concentrates on the diagnosis of tumours in general, it will be appreciated by the skilled person that the present invention may equally and advantageously be applied to the diagnosis of other disease states characterised by gene expression profiles, e.g. hypo/hyperthyroidism, diabetes, or organ rejection.

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Further, the invention may be used to test plasma samples for leukaemia or other hematopoetic disorders.

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In previous studies carried out by the present inventors, a large degree of heterogeneity in protein expression was observed, particularly in malignant tumours (17, 18). Both qualitative and quantitative differences were found within each tumour group.

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However, the large quantitative variability indicated that identification based on pattern recognition would be difficult. However, the present inventors show herein that it is possible to select a subset of variables which show a characteristic pattern within the group, and thus are useful for prediction of the presence of malignant cells and their initial origin.

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Thus, in a second aspect of the present invention, there is provided a method of creating a collection of diagnostic markers based on protein expression levels for use in classifying disease cells in a given sample, comprising

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(1) obtaining cells from a plurality of samples of a selected disease type;

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(2) disrupting the cells to expose the cellular proteins contained therein;

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(3) separating the cellular proteins according to their size on a two-dimensional electrophoresis gel for each of said plurality of samples or a selected disease type; and

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(4) scanning said two-dimensional electrophoresis gels to collect image data for each of the plurality of

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samples of a selected disease type;

(5) analysing said image data in order to identify  
one or more markers characteristic of said selected  
disease type.

5 In an alternative second aspect of the present  
invention, there is provided a method of creating a  
collection of diagnostic markers based on nucleic acid  
expression levels for use in classifying disease cells in  
20 a given sample, comprising

10 (1) obtaining cells from a plurality of samples of a  
selected disease type

25 (2) disrupting the cells to obtain the expressed  
nucleic acid sequences contained therein,

30 (3) separating the expressed nucleic acids sequence  
15 according to their nucleotide sequence using micro-array  
technology for each of said plurality of samples of a  
selected disease type;

35 (4) scanning said micro-array to collect image data  
for each of the plurality of samples of a selected  
20 disease type; and

40 (5) analysing said image data in order to identify  
one or more markers characteristic of said selected  
disease type.

45 Again, the disease type is preferably cancer,  
25 wherein a plurality of samples may be collected from  
tumours of a particular cancer, e.g. ovarian, breast,  
50 skin etc, and its gene expression profile characterised

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10 by the present invention.

15 It is important that the scanning of the  
electrophoresis gel or the micro-array easily identifies  
the separated proteins or nucleic acids respectively.

20 5 Therefore, the method may further comprise the step of  
labelling the obtained proteins or expressed nucleic  
acids. Nucleic acid sequences may be labelled by  
standard techniques known to the skilled person such as

25 10 fluorescent, enzyme or radio-active labelling. As an  
alternative to labelling obtained proteins, the gels may  
be stained with, for example silver nitrate, and scanned  
25 using a laser densitometer. Alternatively, the gels may  
be analysed using computer-assisted microscope to  
facilitate classification. The data obtained and

30 15 statistical comparison may be performed. In particular,  
this is preferably a multivariate characterisation of one  
or more numerical parameters associated with the  
35 proteins. In other words, multivariate analysis of a  
plurality of variables generated by, for example,

40 20 computer-assisted image analysis may be easily  
classified. The statistical comparison may, for example,  
identify a sub-set of proteins, from among all of the  
45 proteins on the 2-DE, having a statistically significant

50 25 degree of expression and/or correlation when compared to  
other samples from similar tumour cells. This sub-set of  
proteins may then be used as diagnostic markers for the  
particular tumour or stage of malignancy. Preferably, a

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10 plurality of 2-DE gels are analysed and the distribution pattern of the proteins are determined. A model may then be set up with a specified number of variables between the tumour cells being analysed. For example, a

15 5 comparison may be made between benign/borderline/malignant. Preferably the number of variables separating the groups whether proteins or expressed nucleic acid sequences, will range between 20 and 500, more preferably 50 and 300, even more preferably 20 100 and 200. In general, it is preferably that the number of variables is at least 20, more preferably at 25 50 and even more preferably at least 70, 100 or 150 variables. In the present case, the inventors used 170 variables.

30 15 Quantification and multivariate characterisation of the expression profiles of selected protein or nucleic acid groups may be performed on image analytical data 35 obtained from analysis of the 2-DE or the micro-array respectively and used for objective classification of the 20 tumour cells in a given sample. The multivariate 40 characterisation may be carried out by partial least squares discriminant analysis (PLS-DA). This process allows (i) the construction and characterisation of a 45 protein or nucleic acid expression profile database and 50 data extraction of a plurality of sets of proteins or nucleic acids which contribute significantly to the diagnosis/classification of a disease state; (ii) add

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samples/protein or nucleic acid expression profiles to the database and further improve the future accuracy of the diagnosis/classification; and (iii) query the database via the expert system using new tumour samples/protein or nucleic acid expression patterns aiming at a prediction of diagnosis.

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A protein expression profile database comprising image data which has been analysed in order to determine a plurality of variables for use as diagnostic markers; said data being obtained from analysis of two-dimensional electrophoresis gels showing characteristic protein distribution associated with a disease type or state of development of said disease for use in disease diagnosis forms another aspect of the present invention.

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30 15 A nucleic acid (mRNA or cDNA) expression profile database comprising image data which has been analysed in order to determine a plurality of variables for use as

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diagnostic markers; said data being obtained from

analysis of a micro-array showing characteristic

20 expressed nucleic acid sequence distribution associated

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with a disease type or stage of development of said

disease, for use in disease diagnosis forms yet another

45 aspect of the present invention.

50 25 In a further aspect, the present invention provides

a method of determining the presence, type or stage of a

disease type in a patient comprising the steps of

55 (1) extracting a sample of candidate disease cells

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from the patient;

(2) disrupting the cells so as to expose the  
10 cellular proteins contained therein;

(3) separating said cellular proteins on a two-  
5 dimensional electrophoresis gel; and

(4) analysing said gel by computer assisted image  
15 evaluation so as to compare protein distribution on gel  
with a database of diagnostic markers characteristic of a  
20 plurality of disease types or stages of disease

10 development to determine presence, type or risk of said  
disease in said patient.

25 The present invention also provides a method of  
determining the presence, type or stage of a disease in a  
patient comprising the steps of

30 15 (1) extracting a sample of candidate disease cells  
from a patient;

35 (2) disrupting the cells so as to obtain the  
expressed nucleic acid sequences contained therein;

40 20 (3) separating said expressed nucleic acid sequences  
on a micro-array according to their nucleotide sequence;

45 and

50 25 (4) analysing said gel by computer assisted image  
evaluation so as to compare expressed nucleic acid  
distribution on said micro-array with a database of  
diagnostic markers characteristic of a plurality of  
disease types or stages of disease development to  
determine presence, type or risk of said disease in said

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patient.

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Preferably, the disease type is cancer and the disease cells are tumour cells.

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Sample preparation may be carried out using standard techniques. One typical sample may contain approximately one million cells. Samples may be collected using fine needles aspiration biopsy (FNA) - a routine technique used for cytological diagnosis. The major advantage of using FNA combined with the expert system is (i) early diagnosis if possible, a prerequisite for making early decisions on therapy (ii) effects of hormone - or chemotherapy can be followed at protein expression level, providing early information on e.g. resistance against treatment; and (iii) the analysis is based on an average expression profile of the cell population.

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Samples may also be collected after surgery for analysis in order to guide pathological examination and selection of post-operation therapeutic strategy.

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As mentioned above, the earliest stages of malignant

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tumours are hard to identify and pathologists are rarely

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sure how or where a malignancy began. The present

invention therefore has further utility in being able to

more accurately determine the primary origin of tumour

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cells as the primary tumour and its corresponding

metastasis express very similar 2-DE protein profiles

(Franzen et al, Int. J. Cancer 1996, 69, 408-414). Such

analysis will therefore assist a clinician in determining

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the location of the primary tumour.

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The above disclosure concentrates on the analysis and diagnosis of tumours. However, as mentioned above, the present invention may also be usefully applied to the diagnosis of any disease state that can be characterised by a statistically significant protein expression profile which allows the identification of specific diagnostic markers.

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By way of example only, a brief outline/workflow on how the computer analysis may be set up in practice is provided below:

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1. A new tumour sample is prepared, analyzed by 2-DE and the expression pattern is scanned.
- 30 15 2. All protein spots in this expression pattern is quantified and matched against a reference pattern using any established software for basic 2-DE analysis (e.g. PDQuest, Melanie, BioImage).
- 35 20 3. The data is first organized in a Excel-spreadsheet-like format table with all protein spot reference numbers in the first column and individual normalized protein quantities for every analyzed sample in the following columns. A new case/pattern is added as a new column. This corresponds to the "data table X".

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4. The process of "data mining" - to find those proteins/variables which contribute most to the separation of tumour classes - and build the learning set (the core of the database), is based on the PLS-DA analysis. Here, an additional "data table Y" is included, as described under materials and methods, data preprocessing (please see also references 14 & 15). Graphically and numerically it is possible to make a first selection of variables (those that are far from origo (compare fig. 4) in the same and opposite direction from the corresponding position of tumour classes, compare fig. 3).

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15 5. In an interactive sub-routine or process, this first set of variables is crossvalidated by excluding cases one by one in sequences, rebuild the model and make a prediction of each of the excluded cases.

20 Then, a second set of variables are selected (according to step 4), and so on - until the predictive value reach an optimum. In the present case, a set of 170 variables was selected in this way (step 4 and 5) and is therefore not a random choice.

25 6. Next, the true predictive value is determined using a new set of cases (the test set).

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7. This process, step 3-6, can then be repeated with an increased number of cases in order to further improve the predictive accuracy.

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5 8. A new case (an unknown tumour sample) is then analyzed by 2-DE/basic image analysis, the pattern is compared with respect to the defined group of variables in the database model and classified using, for example, PLS-DA prediction in order to obtain a diagnosis. Each new case may also be added to the database for future improvements of the predictive value of the model.

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15 One part of the expert system/computer software is to integrate steps 3 to 7 and make the process user-friendly in order to guide the investigator towards the construction of a model within the data base which provide high predictive accuracy. The other part of the expert system/computer software is to facilitate the 20 query of the model using a new case in order to obtain a diagnosis (step 8 above). In addition to these 40 "calculation parts" of the expert system, information may be included on sample preparation and on sample 45 characteristics, 5-year survival data etc.

25 Thus, in the further aspect of the present invention, there is a provided a diagnostic kit for diagnosing the presence, type or stage of a disease, e.g.

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a tumour or malignancy of a tumour, said kit comprising a database capable of quantifying an protein or nucleic acid expression pattern and comparing it against reference patterns held within the database. The kit may 5 also optionally include, instructions for carrying out any of the methods described above; apparatus for carrying out a 2-DE; micro-array technology or a laser densitometer or other image scanning device.

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Aspects and embodiments of the present invention 10 will now be illustrated, by way of example, with reference to the accompanying figures. Further aspects 25 and embodiments will be apparent to those skilled in the art. All documents mentioned in this text are incorporated herein by reference.

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#### Brief Description of the Drawings

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Fig. 1 The two first principal components scores (t<sub>2</sub> against t<sub>1</sub>) of the 2-DE training data-set (22 gels and 1553 spots). A = benign ovary tumour sample (open 20 circles), B = borderline ovary tumour sample (mixed circles), and C = malignant ovary tumour sample (filled 40 circles).

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Fig. 2 The two first principal components scores (t<sub>2</sub> against t<sub>1</sub>) of the most informative part of the 2-DE 45 training data-set (22 gels and 170 spots). For 25 descriptions, see Fig 1.

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Fig. 3 The two first PLS-DA scores (tPS<sub>2</sub> against

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tPS,) of the entire 2-DE data (40 gels and 170 spots). The samples in the learning-set are indicated using circles (A = benign ovary tumour sample (open circles), B = borderline ovary tumour sample (mixed circles), and C = malignant ovary tumour sample (filled circles). The samples in the test-set are indicated using filled/mixed and open squares in analogy with the learning-set.

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Fig. 4 The corresponding loading plot to Fig. 3 (wc<sub>2</sub> against wc<sub>1</sub>). Indicated are the loading scores for the most significant spots for separation of the three tumour classes.

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Fig. 5 The two first principal components scores (t<sub>2</sub> against t<sub>1</sub>) of breast tumour samples (33 gels and 170 spots). Cases classified as carcinoma are labelled "C" and have filled symbols; cases classified as fibroadenoma are marked with FA and have open symbols.

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#### Detailed Description

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#### 20 1) MATERIALS AND METHODS

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##### Tumour tissue samples

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All samples were obtained within 40 min after resection and tumour cells were enriched as previously described (10). Histopathological characterization was carried out using hematoxylin-eosin stained sections of formalin fixed and paraffin embedded specimens. Tumours

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- 22 -

were classified using the WHO system.

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#### Electrophoresis, scanning and image analysis

2-DE was performed as previously described (11). Resolyte (2%, pH 4 - 8, BDH) were used for isoelectric focussing, 10 - 13% linear gradient SDS-polyacrylamide gels were used in the second dimension. Gels were stained with silver nitrate as described by Rabilloud et al. (12) and scanned at 100 mm resolution using a Molecular Dynamics laser densitometer. Data was analysed using PDQUEST<sup>TM</sup> software (7) obtained from Pharmacia Biotech (Uppsala, Sweden).

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#### Data preprocessing

15 The data from the matchset was exported from PDQUEST gel analysis package in the form of tables, with rows representing gels and columns representing spots (data table X - see references 14 and 15). Before the analysis, the data was standardized by dividing each variable (table 20 column) by its standard deviation, thereby giving each variable the same influence in the analysis. Thereafter the data is centred by subtracting from each column its 40 average.

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#### Data analysis

50 The preprocessed data table (data table X) was analysed by two data analysis methods. The first one,

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Principal Component Analysis (PCA), extracts the information in the data, in form of eigenvectors or principal components. Visually, one can see this as a cloud of points (the individuals cases/gels) in a multidimensional space (each axis's representing each spot). PCA first centers the data. Secondly, it rotates the data in such a way that the greatest amount of linear variation is described by the first component axis, the residual variation is described by the second component axis, and so on. Most of the information is often compressed into two or three components. A more detailed description of PCA may be found elsewhere (13).

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The second data analysis method, Partial Least Squarest (PLS) - Discriminant analysis, was used to classify the cases into the three tumour-classes (benign, borderline or malignant). An additional data table (data table Y) with the classification of the tumours is included into the analysis. Table Y consists of the same number of columns as the number of tumour classes and the number of rows is equal to the number of cases. The table is then filled with suitable dummy variables (i.e. 1 = belongs to a specific tumour class or 0 = does not belong).

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The PLS-analysis is similar to PCA in that it projects the data table X into a vector. It differs, however, in that the direction of the vector is determined both by the variation of data table X (as in the case of PCA) as well as the variation of data table Y. For further descriptions

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of PLS, see (14, 15). The significance of the PLS-model is checked by cross-validation. Data from a small number of samples is kept out of the calculation, the PLS model is computed from the remaining data, and the y-values of the deleted are thereafter predicted from the model. The differences in square between predicted and actual y-values for deleted samples are summed to form PRESS (Predictive Error of Sum Squares). This sequence is repeated until each sample has been deleted once.

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10 The data-table used for training the PLS-model consists of 22 cases and 170 spots (Table X). To test the 25 model a table (18 cases and 170 spots) with unknown tumour class was used (Table X).

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30 The data analysis were carried out on CODEX™ software 15 obtained from Sumit System AB (Stockholm, Sweden) and SIMCA™ software obtained from Umetri AB (Umeå, Sweden).

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## 2) RESULTS

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### Creation of a Learning Set

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40 Cells were extracted from fresh ovarian tumour tissue and single cell suspensions free of erythrocytes were prepared (11). Cytological smears were prepared from all 45 preparations and samples usually contained > 90% tumour 25 cells (histopathological characteristics are presented in Table 1). 2-DE polypeptide patterns obtained from these 50 cells were analysed by the PDQUEST™ software (7). The

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patterns of polypeptide expression in 22 ovarian tumours were examined, 5 benign (A), 6 borderline (B) and 11 malignant (C) cases (objects). These patterns were matched together and a reference 2-DE map was constructed containing 1553 spots (variables).

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As an initial step, principal component analysis was applied to entire material (22 gels and 1553 spots) to provide an overview over the data structure, to identify outliers and possible clusters. Normalized quantities (expressed as ppm) for all spots were used for the PCA.

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Fig. 1 shows the scores for the first two components. A coarse separation into two major groups, A + B and C was observed, indicating that latent structures with predictive value are present in this set of data. However, the corresponding loading plots showed very scattered data (data not shown).

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Of the original data (1553 variables, Fig. 1), 170 variables had a substantial influence on the model (PLS loadings  $> 0.02$ ). Approx. 100 variables were active in separating the groups A + B (benign/borderline) and C (malignant), and approximately 70 variables in separating between A (benign) and B (borderline). An improved separation of the clusters representing each of the three classes was observed using these 170 variables (Fig. 2).

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Four significant PLS-DA vectors were found, by using cross-validation ( $Q^2=0.84$ ), describing 98.4 % of the variance in Y and 40.7 % in X. This data set was then closed and

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10 called "learning set".

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Testing the model with unknown tumours

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15 Eighteen new cases were analysed by 2-DE and added to the existing matchset. Expression levels of the 170 markers for all cases were analysed blindly using PCA, enabling the distribution of new objects. Figure 3 show the predictions of unknown cases in a PLS score plot (and the corresponding loadings in Fig. 4).

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20 After breaking the code, 6 of 8 malignant cases were correctly classified. Case 84 and 89 were classified as borderline. Furthermore, 3 of 4 borderline cases were correctly classified, whereas borderline case 96 was classified as benign. Benign cases 90 and 95 were correctly 25 classified. Of the remaining 4 cases, 3 were classified as borderline and one (case 29) as borderline/malignant.

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Testing a ovary model with breast tumours

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20 The possibility that an ovarian cancer model could be used for classification intraductal breast tumours was exploited. The present inventors matched the ovary tumour matchset standard 2-DE map with a corresponding breast tumour standard map in the database (16). Seventy-five of 45 25 the 170 markers were present in the breast standard map.

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Fig. 5 shows the PCA distribution of 33 cases of breast cancer (26 carcinomas, 6 fibroadenomas and 1 normal breast

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epithelium). Only a tendency of clustering of benign cases was observed which indicate that some but not all of the markers show predictive value.

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5 3) DISCUSSION

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The present inventors present here a first attempt to apply artificial learning strategies using quantitative 2-dimensional electrophoresis data for tumour diagnosis. A subset of the information in the 2-DE pattern, based on 170 spots, was selected. Using these variables, a learning set was constructed where an acceptable separation of the groups benign/borderline/malignant tumours into three clusters was obtained. Whether other combinations of spots will result in an improved separation is unknown and difficult to test, since each learning set has to be tested by a new panel of unknown samples. We tested the learning set using 18 cases, and observed a correct classification of the majority of these (11/18).

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It is well known among pathologists that the routinely used limited number of diagnostic sections may not be representative for a certain lesion. In this context it is important to note that the sampling technique employed for 2-DE analysis is more likely to meet the requirements for lesion representivity.

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In previous studies by the present inventors, a large degree of heterogeneity in polypeptide expression was observed, particularly in malignant tumors (17, 18). Both

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qualitative and quantitative differences were found within each tumour group. Particularly, the large quantitative variability indicated that identification based on pattern recognition would be difficult. The present data suggests that it is possible to select a subset of variables which show limited variability within the group, and useful for prediction.

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Neural networks and artificial learning has been used to predict cancer prognosis and for grading tumors (5, 19-22). The parameters used have been various TNM-scoring systems, nuclear grading, tumour markers and histopathological scoring. For prostate cancer, the sensitivity of the network was between 81 to 100% and the specificity 72 to 75% to predict various outcomes such as seminal vesicle and lymph node involvement (22). Similarly, neural network analysis has been performed on breast cancer, using parameters such as hormone receptor status, DNA index, tumour size, number of axillary lymph nodes involved with tumour as input information (20). These studies have indicated that artificial learning is a powerful method to increase the diagnostic accuracy on individual tumours.

The present inventors have noted that many of the alterations observed in 2-DE pattern are similar between tumours of epithelial origin. Thus similar changes in the expression of some cytoskeletal and stress proteins are observed in breast, ovarian and prostate tumors (10; Alaiya

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et al., unpublished). With this background, it was interesting to examine whether a selected set of ovarian markers could be used for classification of intraductal breast tumors into benign and malignant. Some clustering of benign cases was observed, whereas malignant cases showed extensive scattering. It seems reasonable to suggest that it will be difficult to construct a universal model for epithelial tumors, and that learning sets have to be created for each tumour type.

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10 In conclusion, the present study suggests that artificial learning strategies can be used for tumour diagnosis.

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**Claims**

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## Claims

1. A method of obtaining combinations of gene expression profiles in order to determine diagnostic markers characteristic of a selected disease type or stage of development of a disease comprising

(1) obtaining cells from a sample of said disease tissue;

(2) disrupting cells to expose the cellular products characteristic of gene expression;

(3) separating said cellular products according to their characteristic properties on a substrate; and

(4) carrying out computer-assisted multivariate analysis of the substrate to quantify and characterise the cellular product distribution on the substrate to identify specific diagnostic markers characteristic of said disease.

2. A method according to claim 1 wherein the cellular products characteristic of gene expression are proteins.

3. A method according to claim 1 or claim 2 wherein the substrate is an electrophoresis gel which allows separation of the cellular products characteristic of gene expression according to their size.

4. A method according to claim 3 wherein said gel is 2D-electrophoresis gel.

5. A method according to claim 1 wherein the cellular products characteristic of gene expression are nucleic acid sequences.

6. A method according to claim 5 wherein the nucleic acid sequences are mRNA.

7. A method according to claim 1, claim 5 or claim 6

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wherein the substrate comprises a plurality of binding members capable of binding said cellular products characteristic of gene expression.

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5 8. A method according to claim 7 wherein said binding members are oligonucleotides capable of binding said cellular products characteristic of gene expression according to their nucleotide sequence.

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10 9. A method according to claim 1 or claim 2 wherein said binding members are antibodies.

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10. A method according to any one of claims 7 to 9 wherein is said substrate is a micro-array.

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25 11. A method according to any one of the preceding claims wherein said cellular products characteristic of gene expression are labelled to assist computer-assisted multivariate analysis.

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30 12. A method according to any one of the preceding claims wherein said multivariate analysis is carried out by partial least squares discriminant analysis (PLS-DA).

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25 13. A method according to any one of the preceding claims wherein the disease is cancer and the cells are tumour cells or normal reference cells within a given tumour.

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30 14. A method of creating a collection of diagnostic markers based on protein expression levels for use in classifying disease cells in a given sample, comprising  
45 (1) obtaining cells from a plurality of samples of a selected disease;  
35 (2) disrupting the cells to expose the cellular proteins contained therein;  
50 (3) separating the cellular proteins on a two-

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dimensional electrophoresis gel for each of said plurality of samples of the selected disease; and

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(4) scanning said two dimensional electrophoresis gels to collect image data for each of the plurality of samples of the selected disease.

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15. A method of creating a collection of diagnostic markers based on nucleic acid expression levels for use in classifying disease cells in a given sample,

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comprising

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(1) obtaining cells from a plurality of samples of a selected disease;

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(2) disrupting the cells to obtain the expressed nucleic acid sequences contained therein;

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(3) separating the expressed nucleic acid sequences on a micro-array for each of said plurality of samples of the selected disease; and

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(4) scanning said micro-array to collect image data for each of the plurality of samples of the selected disease.

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16. A method according to claim 14 or claim 15 further comprising the step of analysing said image data in order to identify one or more markers characteristic of said selected disease.

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17. A method of determining the presence, type or stage of a disease in a patient comprising the steps of

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(1) extracting a sample of candidate disease cells from the patient;

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(2) disrupting the cells so as to expose the cellular proteins contained therein;

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(3) separating the cellular proteins on a two-dimensional electrophoresis gel; and

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(4) analysing said gel by computer assisted image evaluation so as to compare protein distribution on gel with a database of diagnostic markers characteristic of a

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plurality of tumour types or stages of malignancy to determine presence, type or risk of said disease in said patient.

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5 18. A method of determining the presence, type or stage of a disease in a patient comprising the steps of

(1) extracting a sample of candidate disease cells from the patient;

(2) disrupting the cells so as to obtain the

10 expressed nucleic acid sequences contained therein;

(3) separating the expressed nucleic acid sequences on a micro-array according to their individual nucleotide sequence; and

(4) analysing said micro-array by computer assisted

15 image evaluation so as to compare expressed nucleic acid distribution on said micro-array with a database of diagnostic markers characteristic of a plurality of

20 disease types or stages of development of said disease to determine presence, type or risk of said disease in said patient.

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19. A method according to any one of the preceding claims wherein the number of markers characteristic of said disease type is in the range of 20 to 500.

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25 20. A method according to claim 19 wherein the number of markers characteristic of said disease type is in the range of 50 to 300.

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30 21. A method according to any one of claims 14 to 20 wherein the disease type is selected from the group cancer, hypo/hyperthyroidism, diabetes, organ rejection, 45 and samples for leukaemia or other hematopoetic disorders.

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22. A method according to claim 21 wherein said disease state is cancer and said disease tissue is a tumour.

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5 23. A protein expression profile database comprising  
10 image data which has been analysed in order to determine  
15 a plurality of variables for use as diagnostic markers;  
said data being obtained from analysis of two dimensional  
20 electrophoresis gels showing characteristic protein  
distribution associated with disease type and state of  
disease for use in disease diagnosis.

15 24. A protein expression profile database according to  
10 claim 23 wherein said disease is cancer and the state of  
20 said diseases equates to the state of malignancy of said  
cancer.

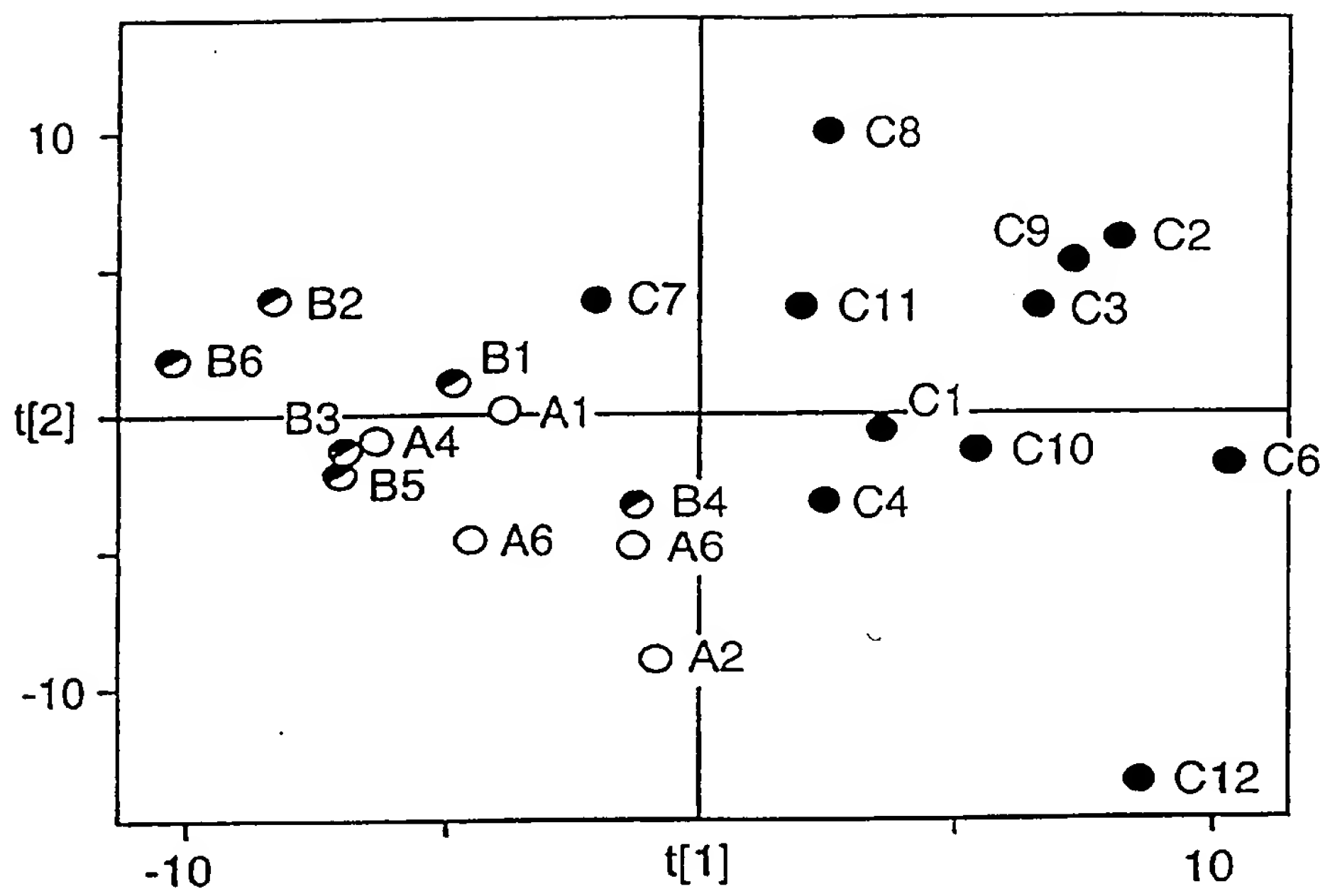
25 25. A nucleic acid expression profile database  
30 comprising image data which has been analysed in order to  
determine a plurality of variables for use as diagnostic  
25 markers; said data being obtained from analysis of a  
micro-array showing characteristic expressed nucleic acid  
30 distribution associated with disease type and state of  
disease for use in disease diagnosis.

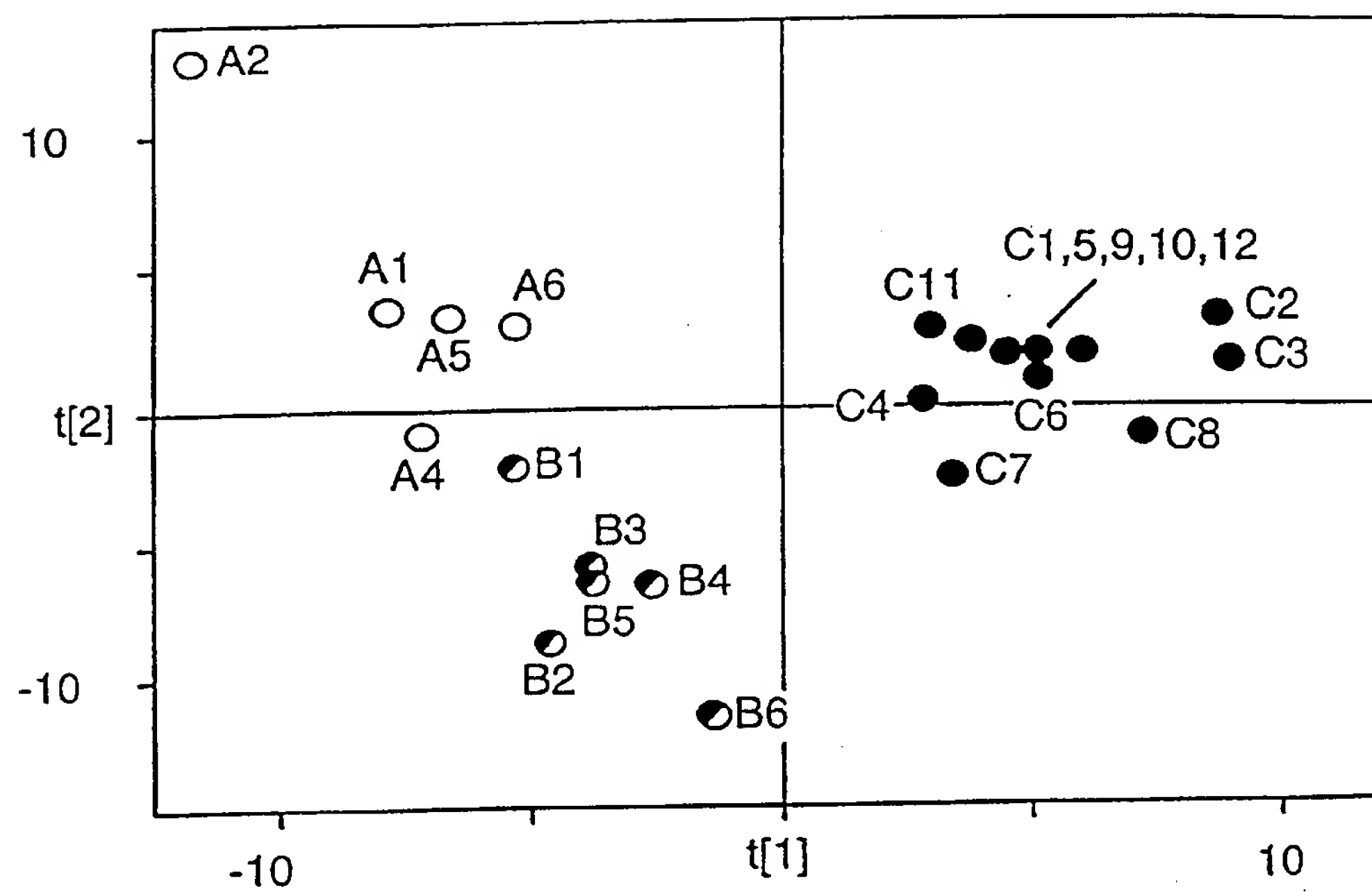
35 26. A nucleic acid expression profile database according  
40 to claim 25 wherein said disease is cancer and the state  
25 of said diseases equates to the state of malignancy of  
said cancer.

45 27. A nucleic acid expression profile database according  
50 to claim 25 or claim 26 wherein the expressed nucleic  
acid is mRNA or cDNA.

Table 1: Histopathological characteristics of samples

Serial No.	Case- No.	Learning model label	Test Cases: Predicted Result from PLS-DA	True Type (A/B/C)	Pathological Diagnosis
1	OC14	A1		A	Serous Cystadenoma IA
2	OC19	A2		A	Serous Cystadenoma IA
3	OC34	A4		A	Serous Cystadenoma IA
4	OC38	A5		A	Serous Cystadenoma IA
5	OC26	A6		A	Mucinous Cystadenoma IIA
6	OC82		B	A	Cystadenofibroma
7	OC39	B1		B	Borderline Seropapillary IB
8	OC46	B2		B	Borderline Seropapillary IB
9	OC50	B3		B	Borderline Seropapillary IB
10	OC21	B4		B	Borderline Mucinous IIB
11	OC59	B5		B	Borderline Mucinous IIB
12	OC68	B6		B	Borderline Mucinous IIB
13	OC72		B	B	Borderline Serous
14	OC77		B	B	Borderline Serous
15	OC07	C1		C	Sero Papillary ADC( IC)
16	OC08	C2		C	Sero Papillary ADC( IC)
17	OC09	C3		C	Sero Papillary ADC( IC)
18	OC20	C4		C	Seropapillary IC
19	OC30	C6		C	Bil Seropapillary IC
20	OC40	C7		C	Bil Adenocarcinoma
21	OC43	C8		C	Bil Seropapillary IC
22	OC04	C9		C	Mixed tumor
23	OC06	C10		C	Clear Cell tumor (IVC)
24	OC27	C11		C	Clear Cell tumor (IVC)
25	OC33	C12		C	Endometrioid Ca IIIC
26	OC48		C	C	Sero Papillary IC
27	OC45		C	C	Endometrioid Ca IIIC
28	OC90		A	A	Serous Cystadenofibroma
29	OC96		A	B	Borderline Serous
30	OC49		C	C	Endometrioid Ca IIIC
31	OC84		B	C	Clear Cell tumor (IVC)
32	OC74		C	C	Endometrioid Ca IIIC
33	OC73		C	C	Sero Papillary ADC( IC)
34	OC89		B/C	C	Sero Papillary ADC( IC)
35	OC95		A	A	Serous Cystadenoma IA
36	OC29		B	A	Mucinous Cystadenoma IIA
37	OC66		B	A	Serous Cystadenoma IA
38	OC35		B	A	Serous Cystadenoma IA
39	OC111		C	C	Sero Papillary ADC( IC)
40	OC117		B	B	Borderline Mucinous IIB





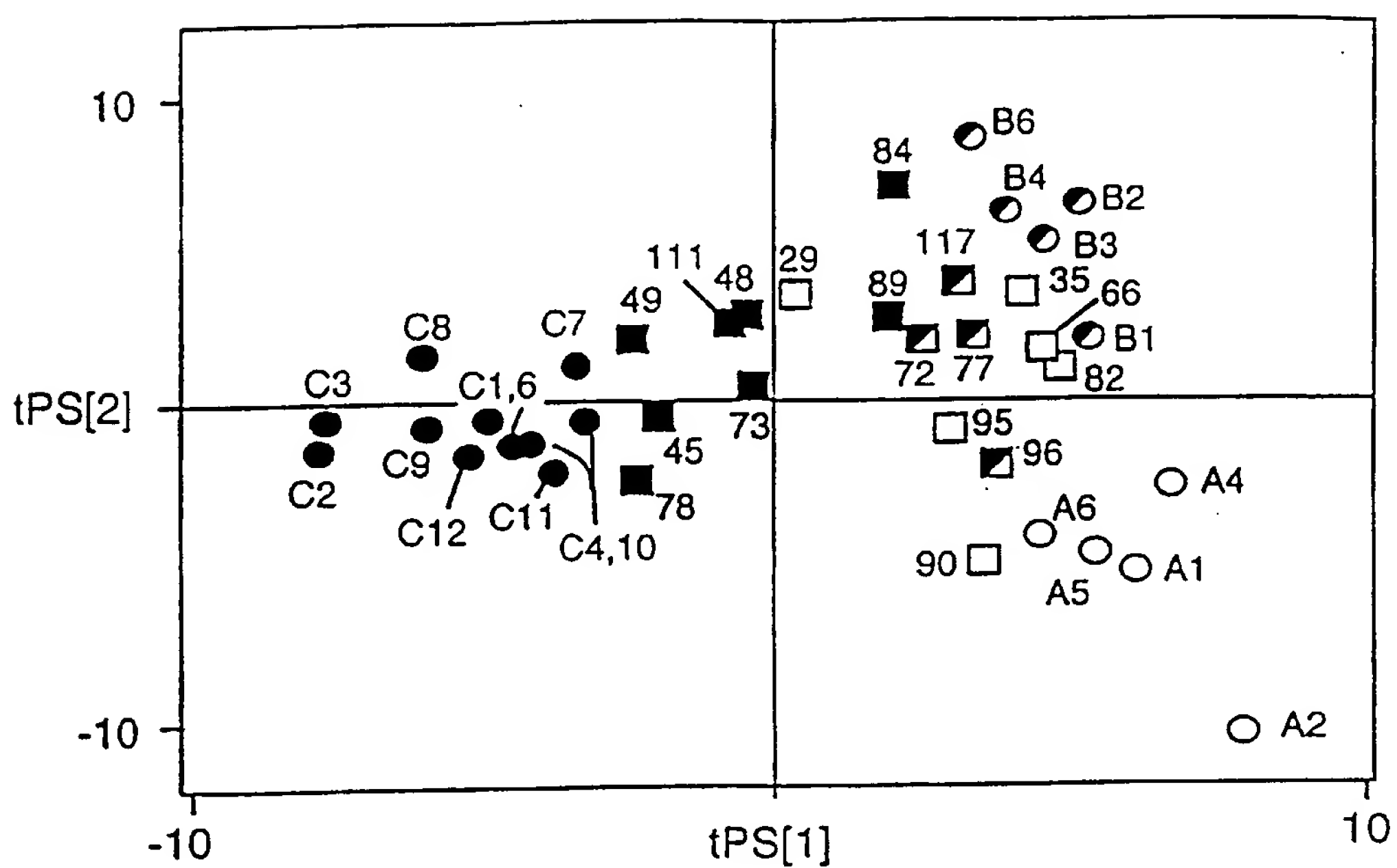


FIG. 3

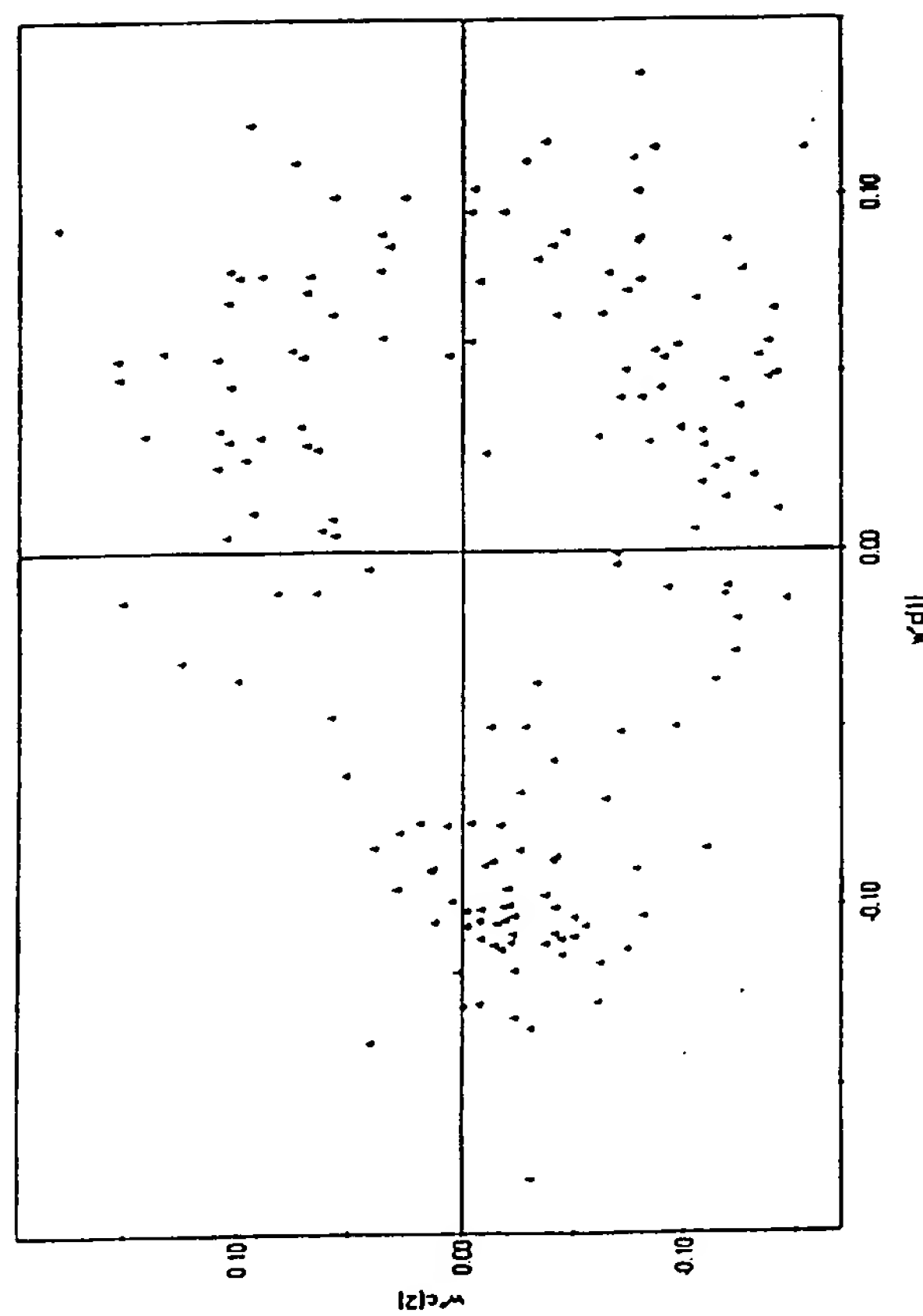


FIG. 4

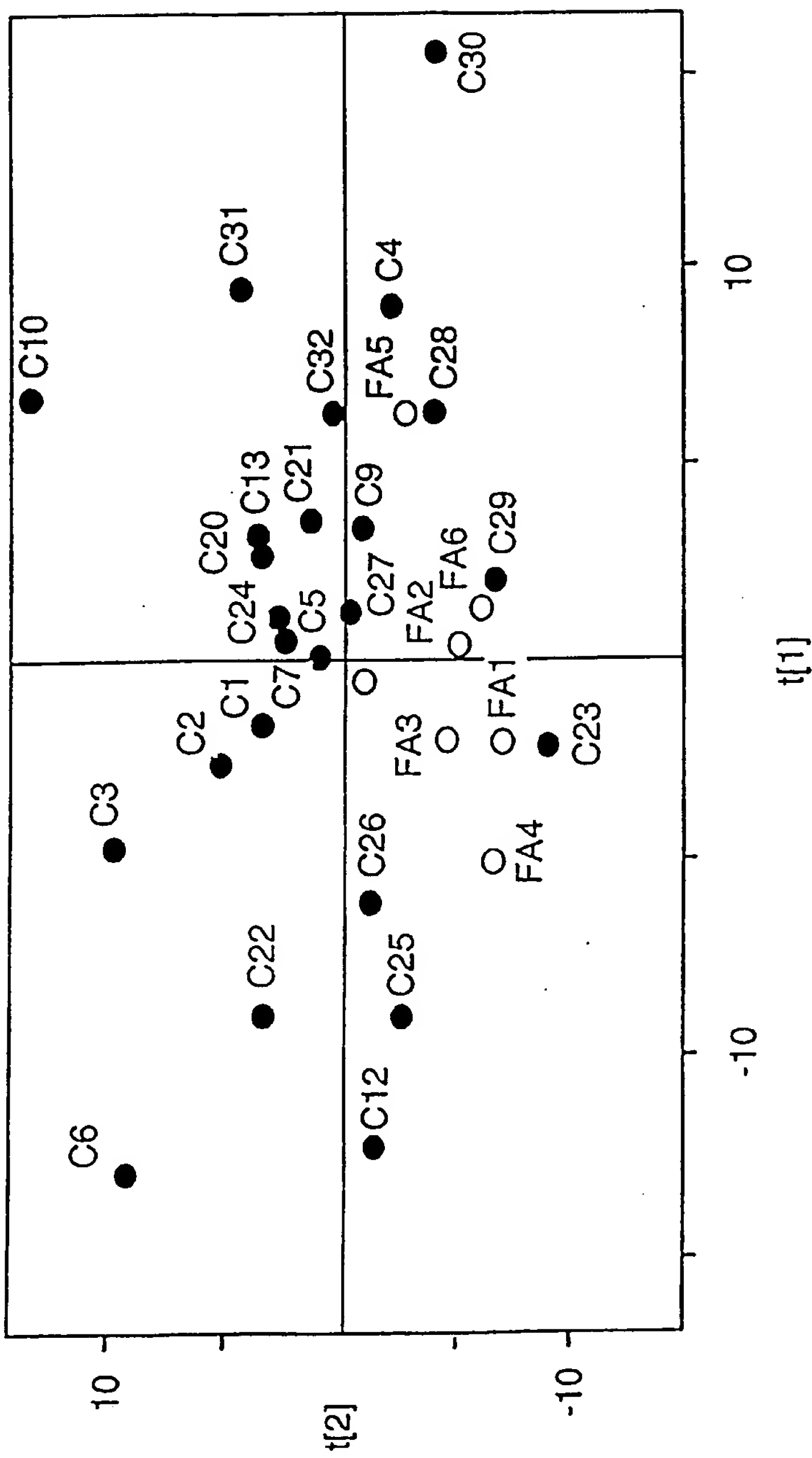


FIG. 5